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1	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 5578175 A	19961126	12	Process for manufacturing iridium and palladium	204/290.12	205/103; 205/206;		Irn, Kwang-lung et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
2	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 5395705 A	19950307		Electrochemical cell having an electrode containing a	429/42	429/30; 429/33;		Door, Robert D. et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
3	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 5296629 A	19940322		Preparation of an electrocatalytic cathode for	502/101	427/115; 429/27;		Marsh, Catherine L. et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
4	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 4514478 A	19850430		Method of making a porous carbon cathode, a porous	429/345	29/623.1; 29/623.5		Binder, Michael et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
5	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 4506028 A	19850319		Process for preparing a fuel cell electrode substrate	502/101	264/29.3; 264/29.5;		Fukuda, Hiroyuki et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

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1	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		US 20020146615	20021010	10	Electrochemical device and method for preparation	429/44	427/115;		Yamaura, Kiyoshi et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
2	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		US 6010606 A	20000104	9	Gas diffusion electrodes	204/284	429/30;		Denton, Jan et al.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
3	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		US 5865968 A	19990202	10	Gas diffusion electrodes	204/284	429/40;		Denton, Jan et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
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US 6010606 A	9			USPAT
US 5865968 A	10			USPAT
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TITLE: Gas diffusion electrodes

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Detailed Description Text - DEX (6):

A first embodiment of the present invention provides a gas diffusion electrode as hereinafter described wherein the catalyst component is one or more metals or their oxides in the form of finely divided unsupported powders or as metals in a dispersed form on a carbon support. Suitably the one or more metals may be a precious metal (Pt, Pd, Ru, Rh, Ir, Os, Au and Ag) or a transition metal selected from groups IVB, VB, VIB, VII, VIII, IX or IIB of the Periodic Table in "Handbook of Chemistry and Physics", 64th Edition, CRC Press, or a combination or alloy thereof. Preferably, the one or more metals is a precious metal, particularly Pt, or an alloy thereof.

Detailed Description Text - DEX (27):

The electrode formed the cathode of an MEA, with the face of the electrode comprising the platinum catalyst component bonded to the membrane electrolyte face. The membrane employed was Du Pont NAFION 112. The single cell results are shown in FIG. 1 and demonstrate that good cell performances were obtained from the MEA comprising the lower cost, more manufacturable electrode of the invention. For operation on pure oxygen very high current densities of over 2.0 A/cm² were obtained. For most practical applications of the PEMFC, the oxidant will be air, and these applications will require that at least a current density of 500 mA/cm² is achieved. As illustrated in the figure, current densities up to 1.0 A/cm² were obtained, and the results represent performances typical of a satisfactorily performing MEA. It is worth noting that on air operation there was a tendency for the cell voltage to decrease more rapidly as the current density increased toward 1.0 A/cm², compared to the pure oxygen data. This is an example of cell voltage decrease due to mass transport losses, relating to the ease with which reactant oxygen in air can diffuse to the electrode reaction sites. This is also a typical characteristic of cell current vs voltage plots seen with conventional MEAs, fabricated with electrodes comprising conducting substrates such as high density carbon fibre paper.

Current US Class - CLASS (2):

429

United States Patent (19)

Denton et al.

Patent Number: 5,865,968
Date of Patent: Feb. 2, 1999

[54] GAS DIFFUSION ELECTRODES

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[73] Assignee: Johnson Matthey Public Limited Company, London, United Kingdom

[21] Appl. No.: 802,556

[22] Filed: Feb. 19, 1997

[30] Foreign Application Priority Data

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Dec. 23, 1996 [GB] United Kingdom 9628802

[51] Int. Cl.⁶ C25B 11/00

[52] U.S. Cl. 204/284; 429/40; 429/41; 429/42

[56] Field of Search 204/283, 284; 429/40, 41, 42

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Primary Examiner—Bruce E. Bell
Attorney, Agent, or Firm—Stevens, Davis, Miller & Mosher, L.L.P.

ABSTRACT

A gas diffusion electrode comprising a non-woven network of fibres, one or more catalyst components and at least one polymeric substance characterized in that the catalyst is embedded within the fibre network is disclosed.

20 Claims, 2 Drawing Sheets

21 East - Detail Last Workspace II In Panel Layout AVE WSP II
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☒ Drafts
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☒ L1: (28595) ("429").CLAS.
☒ L2: (135734) palladium or Pd
☒ L3: (125392) iridium or Ir
☒ L4: (365030) electrode or electrodes
☒ L5: (158983) cathode or cathodes
☒ L6: (23116) 12 same 13
☒ L7: (2552) 16 same 14
☒ L8: (614) 16 same 15
☒ L9: (2858) 17 or 18
☒ L10: (342) 19 and 11
☒ L11: (620876) carbon
☒ L12: (306) 110 and 111
☒ L13: (489) 19 same 111
☒ L14: (120) 113 and 11
☒ L15: (230) high adj density adj carbon
☒ L16: (3) 114 and 115
☒ L17: (2407) carbon adj paper
☒ L18: (23) 114 and 117

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1	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 20030047461	20030313	10	Fuel-cell electrode and method of manufacturing the	205/317	205/109;		Kawahara, Tatsuya et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
2	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 20020132159	20020919		Electrode base material for fuel cell	429/44	429/40;		Ohya, Shyusei et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
3	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 20020098405	20020725		Membrane electrode	429/44	502/182		Odagard, Madeline et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
4	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 20020061431	20020523		assemblies for direct solid polymer electrolyte, a	429/33	429/115;		Koyama, Toru et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
5	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 20020015878	20020207		membrane using thereof, a	429/33	429/314;		Tsumura, Naohiro et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
6	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 20020009630	20020124		Electrodes for fuel cell and processes for producing the	429/42	429/44;		Gao, Yunzhi et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
7	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 20020009630	20020124		Embossed current collector separator for	429/34	502/101		Teraono, Shinji et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
8	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 20020009626	20020124		Polymer electrolyte fuel cell and method for its	429/30	429/42;		Hiromi, Shuji	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
9	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 6538201 B1	20030304	21	Electrode for fuel cell and process for producing the	429/42	429/44;		Medeiros, Maria G. et al.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
10	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 6465124 B1	20021015	5	Magnesium anode, seawater/acid/catholyte	429/105	502/101		Itoh, Takashi et al.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
11	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 6326098 B1	20011204	11	Electrocatalyst, and electrodes,	429/40	429/33;		Narayanan, Sekharipuram R. et al.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
12	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 6171721 B1	20010109	10	Sputter-deposited fuel cell membranes and electrodes	429/41	429/44		Shun, Yoh-Keung et al.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
13	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 6127061 A	20001003	16	Catalytic air cathode for air-metal batteries	429/40	204/192-14;			<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
14	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>					429/133;	204/283;			<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
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US-PAT-NO: 5350643

DOCUMENT-IDENTIFIER: US 5350643 A

TITLE: solid polymer electrolyte type fuel cell

----- KWTIC -----

According to the present invention, the catalytic layers in both the hydrogen electrode and the oxygen electrode comprise a carbon carrier, an active component (catalyst) supported thereon, a proton conductor and a water-repellent binder. The active component is preferably platinum or platinum group metals such as rhodium, ruthenium, palladium and iridium and materials of the proton conductor may be the same as or different from the solid polymer electrolyte. Furthermore, the water repellent binder is preferably a fluorocarbon polymer such as polytetrafluoroethylene (PTFE) or graphite fluoride represented by the formula $(CF)_n$ and/or a mixture thereof.

Detailed Description Text - DETX (19):

An electroded catalyst comprising carbon powders on which platinum was supported was sufficiently kneaded with a perfluorosulfonic acid ion-exchange resin (Nafion liquid manufactured by Alkath Chemical Co.) as a proton conductor and an aqueous suspension of PEPG to prepare a paste. This paste was coated on a carbon paper of about 100 μm in pore diameter and 100 μm thick coated with PEPG which was an electron conductor (gas diffusion layer). This was dried at 80 degrees C. to obtain an electrode. The above electron conductor was obtained by coating an aqueous suspension of PEPG on the carbon paper at a coating amount of 12 mg/cm² and firing it at 350 degrees C. The composition of the hydrogen electrode was as follows: amount of platinum: 0.3 mg/cm², amount of the above proton conductor: 30% by weight and amount of PEPG: 30% by weight. The composition of the oxygen electrode was as follows: amount of platinum: 0.3 mg/cm², amount of the above proton conductor: 20% by weight and amount of PEPG: 20% by weight.

Detailed Description Text - DETX (23):

Electrodes were prepared in the following manner. A catalyst comprising a carbon carrier on which platinum was supported and a petfluorocarboxylic acid resin which was a proton conductor were sufficiently kneaded to obtain a catalyst paste. This paste was rolled by a roll press to obtain a plurality of sheets. These sheets were impregnated with an aqueous PMP suspension having a PMP concentration of 20% by weight and dried at 80 degrees C. to obtain sheet-like catalyst layers. Then, these sheet-like catalyst layers were impregnated with another aqueous PMP suspension having a PMP concentration different from that of the above suspension and dried at 80 degrees C. In this way, electrodes were prepared in which the catalyst layer of both the hydrogen electrode and the oxygen electrode had a concentration gradient of the water-repellant in the thickness direction of the catalyst layer. The catalyst layer of the hydrogen electrode had such a concentration gradient of the water-repellant as 20% by weight in the portion facing to the electrolyte membrane and 40% by weight in the portion facing to the gas diffusion layer. The catalyst layer of the oxygen electrode had such a concentration gradient of the water-repellant as 10% by weight in the portion facing to the electrolyte

Imahashi et al.

[54] SOLID POLYMER ELECTROLYTE TYPE FUEL CELL

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[73] Assignee: Hitachi, Ltd., Tokyo, Japan

[22] Filed: Jun. 1, 1993

Jun. 2, 1992 [JP] Japan

[51]	Int. Cl. ³	H01M 8/10; H01M 4/86
[52]	U.S. Cl.	429/33; 429/30

[58] Field of Search 429/41; 429/42; 429/30, 42, 41, 33

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Primary Examiner—John S. Maples

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ABSTRACT

[57] Provided is a solid polymer electrolyte type fuel cell which is improved in cell output characteristics by preventing the flooding phenomenon at the interface between the oxygen electrode and the electrolyte membrane, accelerating the gas diffusion and effective utilization of the active surface of the catalyst. This solid polymer electrolyte type fuel cell comprises solid polymer electrolyte membrane 1 and gas diffusion electrodes 2 and 3 provided on both sides of the membrane, acid gas diffusion electrodes comprising catalyst layers 6 and 8 and gas diffusion layers 7 and 9 being provided on the outer side of the respective catalyst layers and a hydrogen-containing gas and an oxygen-containing gas being fed to the respective electrodes. In this cell, the water-repellency of the hydrogen electrode 2 is higher than that of the oxygen electrode 3 and furthermore, a gradient of water-repellency is provided in the catalysts in layer of each electrode so that the water-repellency in the portion facing to the electrolyte membrane is higher than that in the portion facing to the gas diffusion layer in each catalyst layer. The output density of the cell according to the present invention is as high as 2-3 times that of conventional cells.

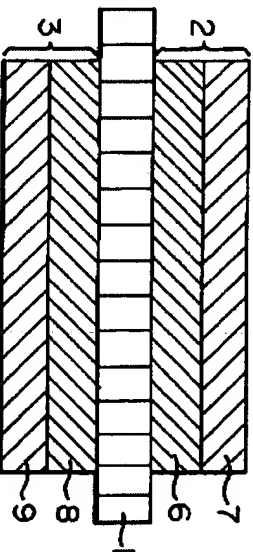
11 Claims, 3 Drawing Sheets

[57] **ABSTRACT**

[57]

Attorneys, Agent, or Firm—Antonelli, Terry, Stout & Kraus

[11]	Patent Number:	5,350,643
[45]	Date of Patent:	Sep. 27, 1994



11 Chains, 3 Drawing Streets

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DOCUMENT-IDENTIFIER: US 6290839 B1

TITLE: Systems for electrophoretic transport and detection of analytes

RWIC

Detailed Description Text - DPMX (342):

In a preferred embodiment, electronic detection is used, including amperometry, voltammetry, capacitance, and impedance. Suitable techniques include, but are not limited to, electrogravimetry; coulometry (including controlled potential coulometry and constant current coulometry); voltammetry (cyclic voltammetry, pulse voltammetry (normal pulse voltammetry, square wave voltammetry, differential pulse techniques); stripping analysis (anodic stripping analysis, cathodic stripping analysis, square wave stripping analysis); conductance measurements (electrolytic conductance, direct analysis); time-dependent electrochemical analyses (chronopotentiometry, amperometry, AC polarography, chronopotentiometry, cyclic voltammetry and amperometry; capacitance measurement; AC voltammetry; and photoelectrochemistry.

Current US Class - CLASS (2):

205

United States Patent

Kayem et al.

SYSTEMS FOR ELECTROPHORETIC TRANSPORT AND DETECTION OF ANALYTES

Inventors: Jon Fuh Kayem, President, Gary Blackburn, President, Stephen D. O'Connor, President, all of CA (US)

Assignee: Clinical Micro Sensors, Inc., Pasadena, CA (US)

Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.

This patent is subject to a terminal disclaimer.

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(22) Filed: Aug. 14, 1998

Related U.S. Application Data

(60) Provisional application No. 60/090,389, filed on Jun. 23, 1998.

(51) Int. Cl.⁷ G01N 27/26

(52) U.S. Cl. 205/771.5; 204/403; 204/450; 204/452; 204/409; 204/600; 204/603

(58) Field of Search 204/403, 413, 204/409, 451, 456, 601, 606, 452, 602, 450, 600, 422/68, 11; 435/6, 183; 205/771.5, 775

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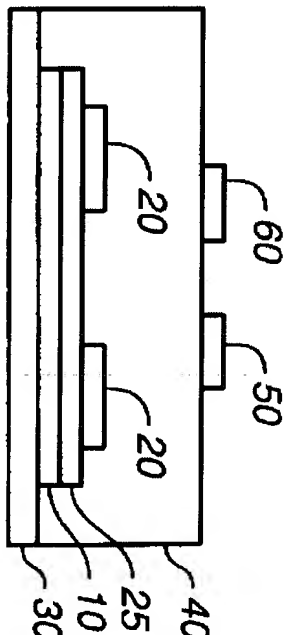
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ABSTRACT

The invention relates to compositions and methods useful in the electrophoretic transport of target analytes to a detection electrode comprising a self-assembled monolayer (SAM). Detection proceeds through the use of an electron transfer moiety (ETM) that is associated with the target analyte, either directly or indirectly, to allow electronic detection of the ETM.

28 Claims, 21 Drawing Sheets



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US 5382331 A	16	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		USPAT
US 5059290 A	3	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		USPAT

US-PAT-NO: 5059290

DOCUMENT-IDENTIFIER: US 5059290 A

TITLE: Electroanalytical method

RWIC -----

Detailed Description Text - DCTX (26):

Using as a working electrolyte, an aqueous solution of acetic acid-sodium acetate containing KI of pH 3.5 and on the other hand, using as a counter electrode solution, an aqueous solution of acetic acid-sodium acetate containing iodine and iodine ion and at a potential of the working electrode of -0.5V vs. the counter electrode, residual chlorine in tap water was determined (controlled potential coulometry).

Detailed Description Text - DCTX (51):

The determination in Example 4 was carried out according to a constant-current method of seeking the endpoint from change in the potential. As a result, the determination could be carried out with an accuracy almost same with that in the case of controlled potential coulometry.

Detailed Description Text - DCTX (62):

The concentration of L-ascorbic acid of reduced type in various foods were measured according to controlled potential coulometry using ferricyan ion as oxidation mediator. The construction of the detector was made as follows:

Current US Class - CLAS (2):

205

United States Patent [19]

Uehyama et al.

(11) Patent Number: 5,059,290

(45) Date of Patent: Oct. 22, 1991

[34] ELECTROANALYTICAL METHOD

[75] Inventors: Shunichi Uehyama, Fukuoka; Shunichi Sasaki, Tokyo, both of Japan

[77] Assignee: Mitsui Engineering & Shipbuilding Co. Ltd., Tokyo, Japan

[21] Appl. No.: 301,792

[22] Filed: Jan. 28, 1989

[30] Foreign Application Priority Data

Jan. 29, 1988 [JP] Japan 63-16895

[31] Int. Cl.³ G01N 27/26

[32] U.S. Cl. 204/133.1; 204/131.12;

[38] Field of Search 204/400, 204/403, 204/409

204/431, 432, 403, 153.1, 153.12

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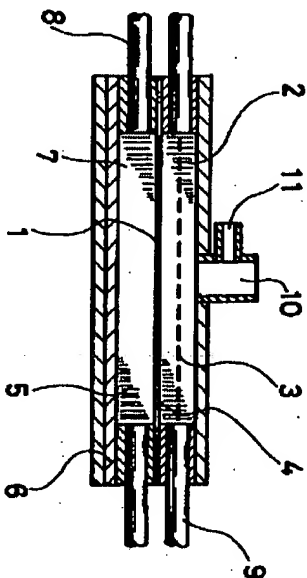
3,924,102 12/1975 Takaue 204/409

Primary Examiner—T. Tung
Attorney, Agent, or Firm—Fey, Sharpe, Beall, Fagan,
Munich & McKee

ABSTRACT

An electroanalytical method which can detect and determine a substance in a short time, with stability and simply is provided, which method comprises providing an electrolytic cell provided with a working electrode chamber and a counter electrode chamber adjacent thereto by the medium of a separator electrolyzing a sample to be determined, by feeding it to a working electrode contained in the working electrode chamber and consisting of an electroconductive porous body impregnated with an electrolyte in a non-flowing state, and measuring at least one of the electric voltage, electric current and electrical quantity in the working electrode, to determine the substance in the sample.

8 Claims, 2 Drawing Sheets



9/2003 09/632,011

3 EAST - Default EAST Workarea (1st Panel/LANDSCAPE) wip.11

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I1: (28887) ("205").CLAS.

I2: (62) controlled adj potential adj coulometry

I3: (20) 11 and 12

I4: (1133) cyclic adj voltammetry

I5: (4) 12 same 14

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DB: USPA/USFORUB

Default operator: OR

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I2: (62) controlled adj potential adj coulometry

I3: (20) 11 and 12

I4: (1133) cyclic adj voltammetry

I5: (4) 12 same 14

I6: (14545) 205/50-333.CCLs.

I7: (0) 12 and 16

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9/2003 09/632,011

2 EAST - [Default EAST WorkSpace (Flat Panel LANDSLAP) web 1]

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L1: (28887) ("205"). CLAS.

L2: (62) controlled adj potential adj coulometry

L3: (20) 11 and 12

L4: (1133) cyclic adj voltammetry

L5: (4) 12 same 14

L6: (14545) 205/50-333. ccls.

L7: (0) 12 and 16

L8: (5399) constant adj potential

L9: (118) 16 and 18

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DB1: USPA/US PUB

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16 and 18

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U	PT	P	Document ID	Issue Date	Pages	Title	Current OR	Current Xref	Retrieval C	Inventor	S	C	3
1			US 20030141193	20030731	16	Methods of anodizing valve metal anodes	205/104	205/148; 205/171		Hossick-Schott, Joachim			
2			US 20030132120	20030717		Method and apparatus for the electrochemical deposition	205/117			Emesh, Ismail et al.			
3			US 20030029728	20030213		Process to separate the vanadium contained in	205/98	205/238; 423/65		Scharifker, Benjamin et al.			
4			US 20020189665	20021219		Preparation of CIGS-based solar cells using a buffered seed layer processes	136/262	136/265; 205/80;		Bhattacharya, Raghu Nath			
5			US 20020134664	20020926		Combinatorial electrochemical deposition	205/118	205/123		Calvert, Jeffrey M. et al.			
6			US 20020100692	20020801		Energy enhanced process for treating a conductive	205/118	204/232; 427/272;		Warren, Christopher J. et al.			
7			US 20020054998	20020509		Catalyzed porous carbon	428/446	204/292; 205/316;		Heimann, Robert L. et al.			
8			US 20020034676	20020321		AN ELECTROACTIVE FILM ON A SUBSTRATE AND METHOD OF	429/44	205/159; 205/224;		Kim, Dong-il et al.			
9			US 20020025449	20020228		Electrolytic system and methods for screening	428/689	205/106; 205/108;		SUKAMTO, JOHANES H. et al.			
10			US 20020014413	20020207		Electrolytic process for treating a conductive	205/81	204/229.8		Symons, Peter G. et al.			
11			US 20020012804	20020131		Patterning of polymer light emitting devices using	428/450	205/106; 205/170;		Heimann, Robert L. et al.			
12			US 6602395 B1	20030805	14		205/317	204/492		Zhuang, Zhiming et al.			

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2 EAST - Default EAST WorkSpace (Last Panel LAMOSTCAPRE.wsp.1)

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Drafts

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Active

L1: (353) (205/102-104).CCUS.

L2: (135734) palladium or Pd

L3: (125392) Iridium or Ir

L4: (23116) 12 same 13

L5: (11) 14 and 11

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U	I	PT	P	Document ID	Issue Date	Pages	Title	Current OR	Current Xref	Retrieval C	Inventor	S	C	3	1
1				US 20030127335	20030710	5	Method for producing material libraries by means	205/82	205/104		Schunk, Stephen A. et al.				
2				US 20020000380	20020103		METHOD, CHEMISTRY, AND APPARATUS FOR NOBLE METAL	205/102	204/212;		GRAHAM, LYNDON W. et al.				
3				US 6350363 B1	20020226		Electric field directed construction of diodes using	205/103	205/114;		Bradley, Jean-Claude				
4				US 6346182 B1	20020212		Process of making bipolar electrodeposited catalysts	205/89	205/102;		Bradley, Jean-Claude				
5				US 6099711 A	20000808		Process for the electrolytic deposition of metal layers	205/101	205/103;		Dehms, Wolfgang et al.				
6				US 5578175 A	19961126		Process for manufacturing Iridium and palladium	204/290.12	205/103;		Lin, Kwang-Lung et al.				
7				US 5495979 A	19960305		Metal-bonded, carbon fiber-reinforced composites	228/124.1	205/103;		Sestri, Surti A. et al.				
8				US 5185073 A	19930209		Method of fabricating dendritic materials	205/104	205/111;		Bindra, Permander S. et al.				
9				US 4909910 A	19900320		Yarns and crows comprising high strength metal coated	205/138	205/104;		Morin, Louis G.				
10				US 4721551 A	19860126		Iridium treatment of neuro-stimulating electrodes	623/24	204/280;		Byers, Charles L. et al.				
11				US 4445980 A	19840501		Copper electroplating procedure	205/103	205/292		Smith, Craig G.				

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	Document ID #	Page	3	X	S	G	P	Kind Codes	Source
1	US 6350363 B1	13	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		USPAT
2	US 6346182 B1	15	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		USPAT
3	US 6099711 A	13	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		USPAT
4	US 5578175 A	12	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		USPAT
5	US 5495379 A	9	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		USPAT
6	US 5185073 A	15	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		USPAT
7	US 4909910 A	8	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>		USPAT

US-PAT-NO: 6346182

DOCUMENT-IDENTIFIER: US 6346182 B1

TITLE: Process of making bipolar electrodeposited catalysts and

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Brief Summary Text - BSTX (15):

One electrodeposition tactic that avoids electrical contact with the conductive support is photoelectrodeposition onto semiconductive particles. This has been used extensively as a method of electrodepositing catalytically active metals (e.g., Au, Pt, Pd, Ag, Ir) onto a dispersed semiconducting support (e.g., TiO₂.sub.2, ZnO, SnO₂.sub.2, ZrO₂.sub.2, CdS, WO₃.sub.3). In this case photons promote electrons from the valence band into the conduction band of the semiconducting particles, creating a situation where anodic and cathodic processes occur on different regions of the same particle. Although this method has been successful in producing catalysts, it requires the use of a semiconductor with a bandgap tuned to the wavelengths capable of penetrating into the sample. Furthermore, due to absorption of light, homogeneous exposures within large volumes is not possible without prolonged mixing.

Detailed Description Text - DETX (14) :

The conductive form of carbon may be graphitic or known forms of conductive diamond. Graphite is preferred. Preferred metals may be Pt, Au, Al, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Ga, Zr, Nb, Mo, Tc, Ru, Rh, Pd, Ag, Cd, In, Sn, W, Re, Os, Ir, Hg, Tl, Pb or Bi and alloys, conductive metal oxides or mixtures thereof. Non-limiting examples of suitable conductive polymers include polypyrrole and its derivatives, polyaniline and its derivatives or polystyrene and its derivatives. A non-limiting example among suitable conductive salt crystal material is a tetrathiafulvene salt. A "conductive semiconductor," as used herein, is one which has sufficient electrically conductive properties to be capable of bipolar polarization using the process of this invention. A conductive semiconductor material preferably comprises doped Si or Ge, or elements of the Periodic Table Groups III and V, such as GaAs, CdSe, InSe, or elements from the Periodic Table Groups II, such as ZnS, CdS, HgSe, ZnS, InSe, ZnTe, HgS, HgTe or HgTe, for example.

Detailed Description Text - DETX (22) :

The source of the catalytic substance is preferably a salt or salts containing an ion of Ed, Pt, Ag, Au, Ni, Cu, Fe, Ru, Rh, Cr, Mn, Ir, Os, Re, Zr, Mo, W, In, Sn, Hg, Tl, Pb, Bi, Cd, Ce and/or mixtures of them. Particularly preferred are salts of Ed, Pt, Fe, Rh, Ag, Au, Co, Fe, and Cu, due to their recognized catalytic properties.

Detailed Description Text - DETX (32):

Preferably the electrodes are comprised of metals or conductive metal oxides of the following group: graphite, Pt, Au, Al, Ti, V, Cr, Mo, Fe, Co, Ni, Cu, Zn, Ga, Zr, Nb, Mo, Te, Ru, Rh, Pd, Ag, Cd, In, Sn, W, Se, Os, Ir, Hg, Pt, Pb, or Bi, or alloys or mixtures of them. Graphite and platinum are the more

United States Patent
(12)
Bradley

Bradley

(10) Patent No.: US 6,346,182 B1
(45) Date of Patent: Feb. 12, 2002

(54) PROCESS OF MAKING BIPOLAR ELECTRODEPOSITED CATALYSTS AND CATALYSTS SO MADE

(75) Inventor: Jean-Claude Bradley, Philadelphia, PA
(US)

(73) Assignee: Drexel University, Philadelphia, PA
(US)

(•) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.

(21) Appl. No.: 09/667,272

(22) Filed: Sep. 22, 2000

Related U.S. Application Data

(63) Continuation of application No. PCT/US99/06430, filed on Mar. 24, 1999.

(60) Provisional application No. 60/079,144, filed on Mar. 24, 1998.

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3661
(a)

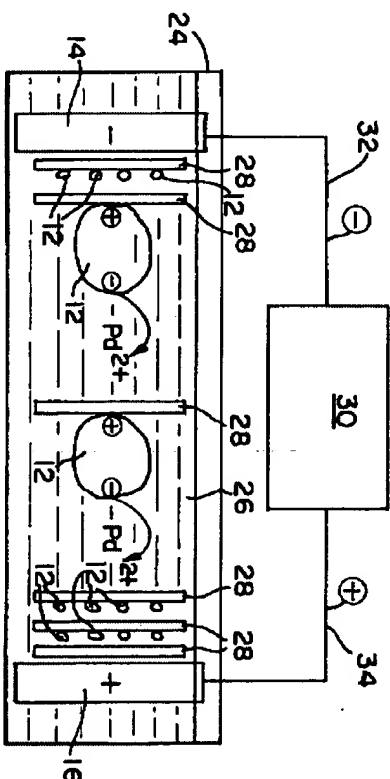
U.S. (52)

(58) Fleet

(56)

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4,031,291 A	6/1977	Fullerwide



25 Claims, 4 Drawing Sheets

1. A process for manufacturing iridium and palladium oxides-coated titanium electrode comprising the steps of:

1. A process for manufacturing an electrode comprising the steps of:

(b) applying iridium and palladium compounds to said titanium substrate to form an iridium and palladium containing layer by a cyclic voltametric deposition process; and

(c) heat-treating said iridium and palladium-applied titanium substrate to obtain an iridium and palladium oxides-coated titanium electrode.

2. A process as claimed in claim 1, wherein said step (b) is executed by immersing said titanium substrate in an iridium and palladium-containing solution to obtain said iridium and palladium containing layer on said titanium substrate by said cyclic voltametric deposition process in said iridium and palladium-containing solution.

3. A process as claimed in claim 2, wherein said iridium and palladium-containing solution comprises a solution of K.sub.2 IrCl.sub.6 PdCl.sub.2, K.sub.2 SO.sub.4 and HCl.

5. A process as claimed in claim 2, wherein said aluminum and aluminum-containing solution has a pH value of about 1.2.

35. An iridium and palladium oxides-applied titanium electrode manufactured by a process as claimed in claim 1, comprising:

(2) an iridium and palladium oxides layer deposited to said titanium substrate.

Current US Cross Reference Classification - CCXR (1)
205/103

Other Reference Publication - ONEP (10):
J. Electroanal. Chem., vol. 256, 1988, pp. 199-205, J. Cox et al.,
"Modification of Glassy Carbon with a Stable Film Containing Iridium Oxide and
Palladium" no month available.

"Electrodes of Conductive Metallic Oxides," Part B. A. Nikola: Technological Impact of Metallic Oxides as Anodes Do data available.

et al., "Microstructure and Electrical Properties of IrO_2 Prepared by Thermal Decomposition of $\text{IrCl}_3 \cdot x \text{H}_2\text{O}$ Role Played by the Conditions of Thermal Treatment" no month available.

Attorney, Agent, or Firm—Oblon, Spivak, McClelland, Meier & Neumann, P.C.

[57]

A process for manufacturing an iridium and palladium oxides-coated titanium electrode comprises preparing a titanium substrate having a surface, applying iridium and palladium to be formed on the surface of the titanium substrate, and heat-treating the iridium and palladium oxides-sprayed titanium substrate to obtain an iridium and palladium oxides-coated titanium electrode. This invention provides a process for obtaining a coated titanium electrode having therein a good adhesion between the coating material and the titanium electrode, and having an excellent electrochemical stability and a superior catalytic activity in an acidic environment.

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- ☐ Drafts
- ☐ Pending
- ☒ Active

- 2011: (353) (205/102-104). CCLS
- 2012: (135734) palladium or Pd
- 2013: (125952) Iridium or Ir
- 2014: (23116) 12 same 13
- 2015: (11) 14 and 11
- 2016: (1033) (205/159-169). CCLS
- 2017: (36) 14 and 16
- 2018: (620876) carbon
- 2019: (22) 17 and 18

Default operator:

☐ **Purest**

☒ **Highlight all hit terms initially**

	U	1	PT	P	Document ID	Issue Date	Pages	Title	Current OR	Current XREF	Retrieval C	Inventor	S	C	3
1	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 20020034676	20020321	18	Catalyzed of fabricating catalyzed porous carbon	429/44	205/159; 205/224;		Kim, Dong-Il et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
2	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 6325910 B1	20011204		Palladium colloid solution and its utilization	205/159	205/125; 205/210		Meyer, Heinrich et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
3	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 6251249 B1	20010626		Precious metal deposition composition and process	205/80	205/159; 205/263;		Chevallier, Jean W. et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
4	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 6051117 A	20000418		Reticulated metal article combining small pores with continuously electroplated foam of improved weight	204/252	204/284; 204/290. 01;		Novak, Donald S. et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
5	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 5804053 A	19980908		Team of improved weight carrier making use of metal catalyst carried on	205/138	204/206; 205/161		Vaccaro, Anthony J. et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
6	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 5531869 A	19960702		Carrier making use of Metal-bonded, carbon fiber-reinforced composites	502/202	205/159; 205/43;		Kubo, Tetsujiro	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
7	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 5495979 A	19960305	9	Process for the metallization of	228/124.1	205/103; 205/161;		Saettri, Surti A. et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
8	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 5421989 A	19950606		Process for the metallization of	205/166	205/159; 205/160;		stamp, Lutz et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
9	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 52556271 A	19931026		Method of immobilizing biofunctional material, and	205/109	204/403.1; 204/403.11;		Ikarizuma, Yoshinoto et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
10	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 5110422 A	19920505		Method for producing an adherent metal deposit on	205/159	205/116; 205/222;		Alperine, Serge et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
11	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 4891069 A	19900102		Composition for the electrolytic coating of	106/1.15	106/1.12; 106/1.13;		Holtzman, Abraham M. et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
12	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	US 4867850 A	19890919		Thermal detectors and process for manufacturing	205/50	205/157; 205/159;		Oka, Syotaro et al.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

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Document ID	Pages	U	V	S	C	P	Kind Codes	Source
4 US 6051117 A	17							USPAT
5 US 5804053 A	13							USPAT
6 US 5531869 A	18							USPAT
7 US 5495979 A	9							USPAT
8 US 5421989 A	11							USPAT
9 US 5256271 A	21							USPAT
10 US 5110422 A	7							USPAT

US-PAT-NO: 5110422

DOCUMENT-IDENTIFIER: US 5110422 A

TITLE: Method for producing an adherent metal deposit on carbon, and mirror obtained by this method

KWIC -----

Abstract Text - ABSTX (1):

A surface layer of a material containing carbon and/or a carbide is produced on the outer surface of a solid carbon-based substrate (1) by selective application of material, said surface layer adhering strongly to the substrate, having a high specific surface area and having open pores (5) of a depth of at least 1 nm, and a metal material having a strong affinity for carbon, comprising at least one metal chosen from cerium, cobalt, chromium, iron, hafnium, iridium, osmium, palladium, platinum, rhodium, ruthenium, lanthanum, manganese, molybdenum, nickel, silicon, tantalum, thorium, titanium, uranium, and tungsten, is deposited on said surface layer, substantially filling said pores.

TITLE - FI (1):

Method for producing an adherent metal deposit on carbon, and mirror obtained by this method

Brief Summary Text - BSTRX (1):

The invention relates to a method for producing a metal deposit on a solid carbon-based substrate, in particular the production of mirrors of low inertia, more particularly for powerful laser beams.

Brief Summary Text - BSTRX (2):

A carbon-based substrate is here understood to be a substrate formed either from a material comprising at least 50% by weight of uncombined carbon or a composite material formed from a disperse phase and a matrix comprising at least 50% by weight of uncombined carbon.

Brief Summary Text - BSTRX (5):

However, in the experience of the authors of the present invention, the adherence of any metal on graphite is a very difficult problem which the document under consideration provides no means of solving. An excellent adherence of the coating is necessary, on the one hand to enable mechanical machining of said coating without it tearing away and on the other hand so that the mirror withstands variations in temperature and the high temperature gradients due to the incidence of powerful rays, taking account of the difference between the coefficient of thermal expansion of carbon and those of the metals, which is very great even when the grade of graphite is chosen so as to limit it, as proposed in the prior application.

Brief Summary Text - BSTRX (6):

The aim of the invention is to provide a method enabling a metal deposit to

United States Patent [19]

Alperthe et al.

US0501032A
[11] Patent Number: 5,110,422
[45] Date of Patent: May 5, 1992

[34] METHOD FOR PRODUCING AN ADHERENT METAL DEPOSIT ON CARBON, AND MIRROR OBTAINED BY THIS METHOD

Inventors:

Serge Alperthe, Paris; Pierre Jossan, Lury Les Moulineaux, both of France

[73] Assignee:

Office National D'Etudes et de Recherches Aeronautiques, Bagneux, France

[21] Appl. No.:

634,593

[22] Filed:

Dec. 11, 1990

[30] Foreign Application Priority Data

Dec. 11, 1989 [FR] France

[51] Int. Cl.

B22D 7/08

[52] U.S. Cl.

427/204; 427/162; 359/900; 359/883; 204/116; 427/204; 427/162; 359/900; 359/883; 204/116; 427/228; 305; 406; 162; 419; 204/15; 38; 1

[56] Field of Search

359/809; 610; 641

[56] References Cited

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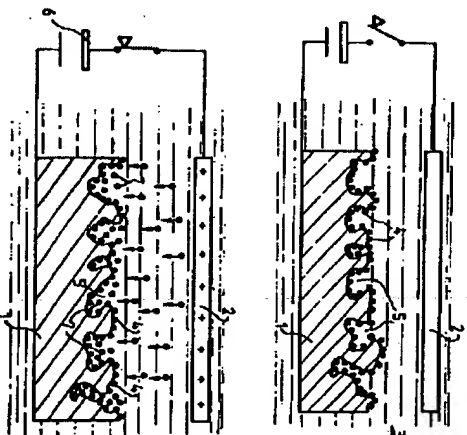
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15 Claims, 2 Drawing Sheets

OTHER PUBLICATIONS

Simon et al., American Ceramic Society Bulletin, vol. 61, No. 8, Aug. 1982, pp. 943-946.

Baudrand, Plating and Surface Finishing, vol. 71, No. 10, Oct. 1984, pp. 72-75.

Primary Examiner—John Niebling

Assistant Examiner—Kishor Mayekar

Attorney, Agent, or Firm—Armstrong, Nikaido, Marmelstein, Kubovick & Murry

ABSTRACT

A surface layer of a material containing carbon and/or a carbide is produced on the outer surface of a solid carbon-based substrate (1) by selective application of material, said surface layer adhering strongly to the substrate, having a high specific surface area and having open pores (5) of a depth of at least 1 nm, and a metal material having a strong affinity for carbon, comprising at least one metal chosen from cerium, cobalt, chromium, iron, hafnium, iridium, osmium, palladium, platinum, rhodium, ruthenium, lanthanum, manganese, niobium, molybdenum, nickel, silicon, tantalum, thorium, titanium, uranium and tungsten, is deposited on said surface layer, substantially filling said pores.

The metal deposit (6) may be rectified and polished without peeling off, in order to produce a mirror of low inertia.

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US 5946222 A	7						USPAT
US 5863400 A	8						USPAT
US 4923574 A	20						USPAT
US 4911798 A	7						USPAT
US 4536259 A	10						USPAT
US 4486274 A	7						USPAT

US-PAT-NO: 5976344

DOCUMENT-IDENTIFIER: US 5976344 A

TITLE: Composition for electroplating palladium alloys and electroplating process using that composition

----- KWIC -----

Brief Summary Text - BSRX (20):

In order to provide a palladium plating bath which results in stable palladium alloy deposition over a wide range of current densities, the present invention employs a mixed ligand system comprising at least a first ligand to complex the palladium and a second ligand to complex a selected alloying base metal. The alloying base metal is used to harden the palladium deposit for increased wear resistance in connector applications and also to lower the cost in other applications such as corrosion protection or decorative applications. The second ligand is chosen to bring the plating potential of the selected alloying base metal and the plating potential of palladium closer together than they would be in the presence of the first ligand alone. By way of example, the base metal may be at least one of the following: iron (Fe), cobalt (Co), ruthenium (Ru), rhodium (Rh) and iridium (Ir).

Detailed Description Text - DSRX (40):

Changes and modifications in the specifically described embodiments can be carried out. For example, based upon the teaching herein, it would be appreciated that in the various Examples 1-4, other alloying metals could also be used to plate Pd alloys including but not limited to Fe, Ir, Rh and Ru. The plating solution taught herein could also be used in plating applications and processes having low current efficiencies (such as strike baths), low metal concentrations as well as low pH values.

Current US Original Classification - CCOR (1):

205/257

United States Patent [19]

Abyx et al.

[11] Patent Number: 5,976,344
[45] Date of Patent: Nov. 2, 1999[54] COMPOSITION FOR ELECTROPLATING
PALLADIUM ALLOYS AND
ELECTROPLATING PROCESS USING THAT
COMPOSITION[73] Inventors: Joseph Anthony Abyx, Warren, N.J.;
Irith Bogachewsky, Naperville, Ill.;
Heinrich K. Strauchl, Summit, N.J.[72] Assignee: Lucent Technologies Inc., Murray Hill,
N.J.

[21] Appl. No.: 08/974,120

[22] Filed: Nov. 19, 1997

Related U.S. Application Data

[63] Continuation of application No. 08/644,347, May 10, 1996,
abandoned.[31] Int. Cl.⁷ C25D 3/50[52] U.S. Cl. 205/257, 205/265, 106/127,
106/128[56] Field of Search 205/257, 259,
205/265, 106/128, 127

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Primary Examiner—Kishor Mayekar
Attorney, Agent, or Firm—Gibbons, Del Deo, Dolan,
Guttmann & Veechman

[57] ABSTRACT

An aqueous electroplating bath for the electrodeposition of
palladium alloys in a mixed ligand system. A first ligand
operates to form a complex of palladium and a second ligand
functions to form a complex of another metal which brings
the plating potentials of the two metals closer together.
Palladium and the alloying metal thus exist as complexes
with different structures.

8 Claims, No Drawings

	Document ID#	Pages	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54	55	56	57	58	59	60	61	62	63	64	65	66	67	68	69	70	71	72	73	74	75	76	77	78	79	80	81	82	83	84	85	86	87	88	89	90	91	92	93	94	95	96	97	98	99	100	101	102	103	104	105	106	107	108	109	110	111	112	113	114	115	116	117	118	119	120	121	122	123	124	125	126	127	128	129	130	131	132	133	134	135	136	137	138	139	140	141	142	143	144	145	146	147	148	149	150	151	152	153	154	155	156	157	158	159	160	161	162	163	164	165	166	167	168	169	170	171	172	173	174	175	176	177	178	179	180	181	182	183	184	185	186	187	188	189	190	191	192	193	194	195	196	197	198	199	200	201	202	203	204	205	206	207	208	209	210	211	212	213	214	215	216	217	218	219	220	221	222	223	224	225	226	227	228	229	230	231	232	233	234	235	236	237	238	239	240	241	242	243	244	245	246	247	248	249	250	251	252	253	254	255	256	257	258	259	260	261	262	263	264	265	266	267	268	269	270	271	272	273	274	275	276	277	278	279	280	281	282	283	284	285	286	287	288	289	290	291	292	293	294	295	296	297	298	299	300	301	302	303	304	305	306	307	308	309	310	311	312	313	314	315	316	317	318	319	320	321	322	323	324	325	326	327	328	329	330	331	332	333	334	335	336	337	338	339	340	341	342	343	344	345	346	347	348	349	350	351	352	353	354	355	356	357	358	359	360	361	362	363	364	365	366	367	368	369	370	371	372	373	374	375	376	377	378	379	380	381	382	383	384	385	386	387	388	389	390	391	392	393	394	395	396	397	398	399	400	401	402	403	404	405	406	407	408	409	410	411	412	413	414	415	416	417	418	419	420	421	422	423	424	425	426	427	428	429	430	431	432	433	434	435	436	437	438	439	440	441	442	443	444	445	446	447	448	449	450	451	452	453	454	455	456	457	458	459	460	461	462	463	464	465	466	467	468	469	470	471	472	473	474	475	476	477	478	479	480	481	482	483	484	485	486	487	488	489	490	491	492	493	494	495	496	497	498	499	500	501	502	503	504	505	506	507	508	509	510	511	512	513	514	515	516	517	518	519	520	521	522	523	524	525	526	527	528	529	530	531	532	533	534	535	536	537	538	539	540	541	542	543	544	545	546	547	548	549	550	551	552	553	554	555	556	557	558	559	560	561	562	563	564	565	566	567	568	569	570	571	572	573	574	575	576	577	578	579	580	581	582	583	584	585	586	587	588	589	590	591	592	593	594	595	596	597	598	599	600	601	602	603	604	605	606	607	608	609	610	611	612	613	614	615	616	617	618	619	620	621	622	623	624	625	626	627	628	629	630	631	632	633	634	635	636	637	638	639	640	641	642	643	644	645	646	647	648	649	650	651	652	653	654	655	656	657	658	659	660	661	662	663	664	665	666	667	668	669	670	671	672	673	674	675	676	677	678	679	680	681	682	683	684	685	686	687	688	689	690	691	692	693	694	695	696	697	698	699	700	701	702	703	704	705	706	707	708	709	710	711	712	713	714	715	716	717	718	719	720	721	722	723	724	725	726	727	728	729	730	731	732	733	734	735	736	737	738	739	740	741	742	743	744	745	746	747	748	749	750	751	752	753	754	755	756	757	758	759	760	761	762	763	764	765	766	767	768	769	770	771	772	773	774	775	776	777	778	779	780	781	782	783	784	785	786	787	788	789	790	791	792	793	794	795	796	797	798	799	800	801	802	803	804	805	806	807	808	809	810	811	812	813	814	815	816	817	818	819	820	821	822	823	824	825	826	827	828	829	830	831	832	833	834	835	836	837	838	839	840	841	842	843	844	845	846	847	848	849	850	851	852	853	854	855	856	857	858	859	860	861	862	863	864	865	866	867	868	869	870	871	872	873	874	875	876	877	878	879	880	881	882	883	884	885	886	887	888	889	890	891	892	893	894	895	896	897	898	899	900	901	902	903	904	905	906	907	908	909	910	911	912	913	914	915	916	917	918	919	920	921	922	923	924	925	926	927	928	929	930	931	932	933	934	935	936	937	938	939	940	941	942	943	944	945	946	947	948	949	950	951	952	953	954	955	956	957	958	959	960	961	962	963	964	965	966	967	968	969	970	971	972	973	974	975	976	977	978	979	980	981	982	983	984	985	986	987	988	989	990	991	992	993	994	995	996	997	998	999	1000	1001	1002	1003	1004	1005	1006	1007	1008	1009	1010	1011	1012	1013	1014	1015	1016	1017	1018	1019	1020	1021	1022	1023	1024	1025	1026	1027	1028	1029	1030	1031	1032	1033	1034	1035	1036	1037	1038	1039	1040	1041	1042	1043	1044	1045	1046	1047	1048	1049	1050	1051	1052	1053	1054	1055	1056	1057	1058	1059	1060	1061	1062	1063	1064	1065	1066	1067	1068	1069	1070	1071	1072	1073	1074	1075	1076	1077	1078	1079	1080	1081	1082	1083	1084	1085	1086	1087	1088	1089	1090	1091	1092	1093	1094	1095	1096	1097	1098	1099	1100	1101	1102	1103	1104	1105	1106	1107	1108	1109	1110	1111	1112	1113	1114	1115	1116	1117	1118	1119	1120	1121	1122	1123	1124	1125	1126	1127	1128	1129	1130	1131	1132	1133	1134	1135	1136	1137	1138	1139	1140	1141	1142	1143	1144	1145	1146	1147	1148	1149	1150	1151	1152	1153	1154	1155	1156	1157	1158	1159	1160	1161	1162	1163	1164	1165	1166	1167	1168	1169	1170	1171	1172	1173	1174	1175	1176	1177	1178	1179	1180	1181	1182	1183	1184	1185	1186	1187	1188	1189	1190	1191	1192	1193	1194	1195	1196	1197	1198	1199	1200	1201	1202	1203	1204	1205	1206	1207	1208	1209	1210	1211	1212	1213	1214	1215	1216	1217	1218	1219	12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US-PAT-NO: 4911798

DOCUMENT-IDENTIFIER: US 4911798 A

TITLE: Palladium alloy plating process

----- RWIC -----

Detailed Description Text - DETX (8) :

Most preferred is the compound *tris*-(hydroxymethyl) aminomethane both because the electroplating potential for the palladium complex is well removed from the hydrogen evolution potential and the electroplating potential for alloying metals (e.g. nickel and cobalt) is close to the electroplating potential for the palladium complex. Typical alloying metals are silver, copper, nickel, cobalt, gold, chromium, manganese, ruthenium, rhodium, platinum and palladium. Particularly useful are copper, nickel, cobalt and silver. Preferred are alloys comprising at least 10 mole percent palladium, remainder copper, nickel, cobalt and/or silver. Other useful alloys are 40, 60 or 80 mole percent palladium, remainder silver; nickel, cobalt and/or silver. Often, the plated palladium or remainder alloy is at least partially covered with a thin (e.g. between 0.01 and 0.5 micrometer thick) layer of gold to improve corrosion resistance and wear resistance.

Detailed Description Text - DETX (26) :

Part of the palladium in the electroplating solution is replaced by the following metals: silver, copper, nickel, cobalt, gold, chromium, manganese, ruthenium, rhodium, platinum and iridium. Mole percent of palladium replaced by these metals were 10, 20, 30, 40, and 90. Electroplating was carried out at 55 degrees C., 55 degrees C. and room temperature.

Claims Text - CLTX (3) =

3. The process of claim 1 in which the metallic substance comprises, in addition to palladium, at least one metal selected from the group consisting of silver, copper, nickel, cobalt, gold, chromium, manganese, ruthenium, rhodium, platinum and iridium.

Current US Cross Reference Classification - CCXR (3) :

205/257

United States Talcum [19]
Abys et al.

- | | | | |
|------|---------------------------------|--|--------------------|
| [34] | PALLADIUM ALLOY PLATING PROCESS | | |
| [35] | Inventor: | Joseph A. Ayra, Warren, Virginia T. | |
| | | Robert, Sumner, Catherine | |
| | | Wolfebald, Chairman Township | |
| | | Morris County, all of N.J. | |
| [73] | Assignee: | AT&T Bell Laboratories, Murray Hill, N.J. | |
| [21] | Appl. No.: | 288,337 | |
| [22] | Filed: | Dec. 20, 1968 | |
| [31] | Int. Cl.: | CSD 3/26; CSD 3/36 | |
| [32] | U.S. Cl.: | 204/44.3; 204/44.5; 204/47.1 | |
| [38] | Field of Search: | 204/44.3; 204/44.5; 204/47.1; 204/47.4; 44.3; 44.4 | |
| [36] | References Cited | | |
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(11) Patent Number: 4,911,198

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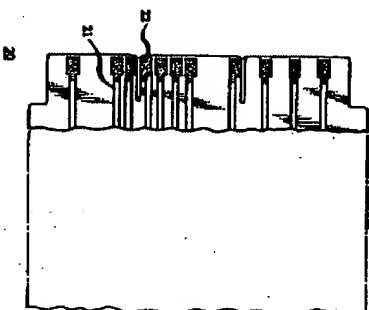
Primary Examiner—G. L. Kaplan
Attorney, Agent, or Firm—Walter G. Nilsen

[57] **ABSTRACT**

ABSTRACT

A process is described for electrophilizing palladium and palladium alloys. The process involves the use of an allyl hydroxyamine as complexing agent and is particularly good for palladium alloys such as palladium-nickel and palladium-cobalt.

11 Chain, 2 Drawing Sheets



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US 4197170 A	9					USPAT
US 4189358 A	10					USPAT

US-PAT-NO: 4189358

DOCUMENT-IDENTIFIER: US 4189358 A

Electrodeposition of ruthenium-iridium alloy

TITLE: KWIC

Brief Summary Text - BSRX (4):

Several baths have been developed for electroplating ruthenium and for electroplating iridium. Examples of ruthenium electroplating baths can be found in U.S. Pat. Nos. 2,057,638, 2,600,175, 3,123,544, 3,576,724, 3,630,856, 3,793,162 and 4,082,625. Examples of iridium plating baths can be found in U.S. Pat. Nos. 1,077,920, 3,554,881, 3,639,219, in Lowenheim's MODERN ELECTROPLATING, 3rd Ed., pp. 354-355 (1974), and in an article by G. A. Corn entitled, "Iridium Plating" in PLATING PROCEDURES, pp. 1258-1261, (1965). In general, ruthenium is considered more difficult to plate than such metals as platinum and palladium, and iridium is considered more difficult to plate than ruthenium. Baths for electrodeposition of certain alloys of ruthenium have also been disclosed, e.g., for Ru-Rh, Ru-Pt, and Ru-Ir in U.S. Pat. No. 3,692,641 and for Rh-Ru in U.S. Pat. No. 3,892,638. None of the patents noted above discloses a bath for co-depositing ruthenium and iridium.

Detailed Description Text - DRTX (2):

FIGS. 1 and 2 are photomicrographs at 500X magnification which show the quality of a Ru-4-6ir alloy deposit from a bath of the present invention on two different surfaces. In both samples the substrate is copper polished metallographically to a 1 μm diamond finish, but in FIG. 1 plating is directly on the copper and in FIG. 2 plating is on copper coated with 0.15 mg/cm. sup. 2 of palladium. FIG. 1, with plating directly on copper, shows cracks at a Ru-Ir loading of 1 mg/cm. sup. 2. FIG. 2, with plating on the palladium coated copper, shows no cracks at a Ru-Ir loading of 1.9 mg/cm. sup. 2.

Detailed Description Text - DRTX (2):

Advantageously, particularly for electroplating applications the valve metal can be coated with a barrier layer, e.g. comprising platinum group metals, gold and nitrides, carbides and silicides of one of the components of the substrate. As shown in FIGS. 1 and 2 a palladium coating, e.g. on a polished copper surface, improved the quality of the deposit. Similar findings have been made with gold and iridium coatings on copper.

Detailed Description Text - DRTX (2):

As used herein, the term "alloy" as applied to a ruthenium-iridium deposit, means that the film contains a mixture of very fine particles of ruthenium and iridium which has a metallic appearance. The particles may be mixed crystals or in solid solution, the microscopic character of the deposited films being different to determine because films are very thin. By "valve" metals is meant those metals form oxide films under anodic conditions, as do, for example, titanium, tantalum, niobium, tungsten, zirconium, aluminum, hafnium and alloys thereof with each other and with other metals. The platinum group metals are platinum, palladium, rhodium, ruthenium, osmium and iridium. The terms electroplated and electrodeposited are used interchangeably. The abbreviations

United States Patent (19)

4,189,358

Scarpellino, Jr et al.

Feb. 19, 1980

[54] ELECTRODEPOSITION OF RUTHENIUM-IRIDIUM ALLOY

[75] Inventors: Anthony J. Scarpellino, Jr, Tucson, N.Y.; William G. Barnes, Ringwood, N.J.

[73] Assignee: The International Nickel Company, Inc., New York, N.Y.

[21] Appl. No. 924,632

[22] Filed: Jul. 14, 1978

[31] Int. Cl. C25D 3/00; C25B 11/00;

[32] U.S. Cl. 204/105 M; 204/106; 204/108; 204/107; 204/109; 204/110; 204/111; 204/112; 204/113; 204/114; 204/115; 204/116; 204/117; 204/123; 204/200R; 204/250 F; 204/293

[38] Field of Search: 204/250 R, 250 F, 293, 204/43 N, 48-49, 112-113, 105 R, 106-108, 109-111, 114-117, 105 M, 123

[56] References Cited

U.S. PATENT DOCUMENTS

3,616,545 10/1971 Bianchi et al. 204/290 F

3,946,573 11/1974 Bianchi et al. 204/250 F

Primary Examiner—R. L. Anderson

Attorney, Agent, or Firm—Evan C. MacQueen, Matthew W. Lott

ABSTRACT

Ruthenium-iridium electrodeposits are prepared from aqueous acid solution containing ruthenium, iridium, a fluoroborate salt, fluoroboric acid, and optionally sulfamic acid. The baths are especially useful for preparing insoluble anodes.

38 Claims, 2 Drawing Figures

9/2003 09632011

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Search

Drafts
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- L1: (353) (205/102-104).CCLS.
- L2: (135734) palladium or Pd
- L3: (125392) Iridium or Ir
- L4: (23116) 12 same 13
- L5: (11) 14 and 11
- L6: (1033) (205/159-169).CCLS.
- L7: (36) 14 and 16
- L8: (620876) carbon
- L9: (22) 17 and 18
- L10: (108) (205/257).CCLS.
- L11: (14) 14 and 110
- L12: (14545) (205/50-333).CCLS.
- L13: (17701) 12 near5 13
- L14: (2460) 113 same 18
- L15: (20) 114 and 112

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114 and 112

PRS Item
BAR Item
Index
Text
HTML

U	PT	P	Document ID	Issue Date	Pages	Title	Current OR	Current Xref	Retrieval C	Inventor	S	C	3
1			US 20030047461	20030313	10	Fuel-cell electrode and method of manufacturing the	205/317	205/109; 429/40;		Kawahara, Tatsuya et al.			
2			US 20020034676	20020321	18	Method of fabricating catalyzed porous carbon	429/44	205/159; 205/224;		Kim, Dong-il et al.			
3			US 20020008038	20020124		Combinatorial Electrochemical synthesis	205/261	204/280; 204/291;		Heller, Adam et al.			
4			US 6280595 B1	20010828		Electrochemical solid phase synthesis	205/122			Montgomery, Donald D.			
5			US 6093302 A	20000725		Electrochemical solid phase synthesis	205/122			Montgomery, Donald D.			
6			US 6051117 A	20000418	17	Reticulated metal article combining small pores with	204/252	204/284; 204/290.01;		Novak, Donald S. et al.			
7			US 5578175 A	19961126	12	Process for manufacturing Iridium and palladium	204/290.12	205/103; 205/206;		Lin, Kwang-Lung et al.			
8			US 5110422 A	19920505	7	Method for producing an adherent metal deposit on	205/159	205/116; 205/222;		Alperine, Serge et al.			
9			US 5085730 A	19920204		Process for regenerating ammoniacal chloride etchants	216/93	204/237; 205/76;		Cordani, John L.			
10			US 4752541 A	19880621		Electrolyte for lithium-sulfur dioxide	429/101	205/59; 252/62.2;		Faulkner, Larry R. et al.			
11			US 4713151 A	19871215		Electrodeposition of lithium	429/325	205/59		Smith, David J.			
12			US 4668347 A	19870526		Anticorrosive coated rectifier metals and their	205/50	205/157; 205/210;		Hebertmann, Clarence E. et al.			

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2 EAST - Default EAST Workpace (Fast Panel LAMDS-CAT) wsp.11

2. EAST - [Default EAST Workspace (Flat Panel LANDSCAPE).wsp:1
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Drafts

- Pending
- Active
 - L1: (353) (205/102-104). CCLS.
 - L2: (135734) palladium or Pd
 - L3: (125392) Iridium or Ir
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 - L5: (11) 14 and 11
 - L6: (1033) (205/159-169). CCLS.
 - L7: (36) 14 and 16
 - L8: (620876) carbon
 - L9: (22) 17 and 18
 - L10: (108) (205/257). CCLS.
 - L11: (14) 14 and 110
 - L12: (14545) (205/50-333). CCLS.
 - L13: (17701) 12 near5 13
 - L14: (2460) 113 same 18
 - L15: (20) 114 and 112
 - L16: (14708) electrodeposits
 - L17: (29486) electroplats
 - L18: (3176) electrochem's near2 deposits
 - L19: (1134) electrochem's near2 coats
 - L20: (41666) 116 or 117 or 118 or 119
 - L21: (275) 113 same 120
 - L22: (164) 121 and 18

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EAST - [Default EAST] Workspace [Flat Path LANDSCAPE.msp:1]

(1) The carbon used as an electrode material readily deteriorates at high temperatures in the presence of both sodium hydroxide and oxygen to considerably impair electrode performance.

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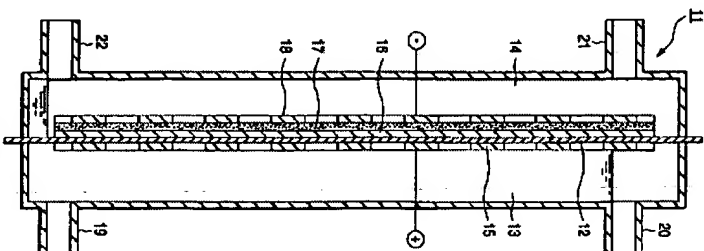
(7) If glass utilized as an oxygen-containing gas, the gas-diffusing ability of the electrode is reduced. This is because carbon dioxide contained in the air reacts with sodium hydroxide to deposit sodium carbonate on the walls of the pores of the gas diffusion electrode.

The oxygen gas diffusion cathode for use in the present invention may have the characteristics of conventional oxygen gas diffusion cathodes. For example, a gauze, sintered powder, sintered metal fiber, formed object, etc., made of a corrosion-resistant material such as, e.g., titanium, niobium, tantalum, stainless steel, nickel, zirconium, gallium, or silver can be used as an electrode substrate, optionally after having been cleaned in a pretreatment. It is preferred to impart moderate porosity and electroconductivity to this electrode substrate so as to smoothly conduct the feeding and removal of electric current and also of the gas and liquid.

A catalyst layer is desirably formed on the surface of the electrode substrate. The catalyst can be made of a metal such as, e.g., platinum, palladium, ruthenium, iridium, copper, silver, cobalt, or lead or an oxide of any of these metals. A layer of the catalyst can be formed by mixing a catalyst material powder with a binder, e.g., a fluorocresin, and a solvent, e.g., naphtha, depositing the resultant paste on the substrate, and solidifying the deposit, or by applying a solution of a salt of a catalyst metal on the substrate surface and burning the coating, or by subjecting the substrate to electroplating in the salt solution or to electrolysis plating in the salt solution in the presence of a reducing agent.

The hydrophilic material interposed between the ion-exchange membrane and the gas diffusion cathode in the present invention is preferably a porous structure comprising a corrosion-resistant metal or resin. This hydrophilic material need not have electroconductivity because it does not contribute to electron movement. Examples of the hydrophilic material include carbon, ceramics such as zirconium oxide and silicon carbide, hydrophilized resins such as PTFE and EBP, metals such as nickel, stainless steel, and silver, and alloys of such metals. The hydrophilic material is preferably in the form of a sheet having a thickness of from 0.01 to 10 mm. Because the hydrophilic material is interposed between the membrane and the cathode, it is desirably an elastic material which, when pressure unevenness is present, deforms to absorb the pressure. Furthermore, the hydrophilic material preferably is made of such a material, and has a structure which can hold a catholyte. Examples of such structures include nets, woven fabrics, nonwoven fabrics and foamed objects. Especially preferred is a sintered plate obtained by mixing a powdery starting material with a pore-forming agent and any of various binders, molding the mixture into a sheet, removing the pore-forming agent with a solvent, and then sintering the sheet, or a structure composed of such sintered plates superposed on each other. An appropriate range of the pore diameter of this hydrophilic

A zero-gg-type electrolytic cell 11 characterized as having hydrophilic liquid-permeable material 16 interspersed between an ion-exchange membrane 12 and a gas diffusion cathode 17. The reaction product passes through the liquid-permeable material and disperses toward edges of the liquid-permeable material before being withdrawn. Hence, the withdrawal direction for the target reaction product is not opposite the feed direction for the reactant gas.



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Document ID	Pages	U	S	C	P	Kind Codes	Source
US 6465124 B1	5						USPAT
US 6436354 B1	15						USPAT
US 6368740 B1	6						USPAT
US 6332900 B1	5						USPAT
US 6326098 B1	11						USPAT
US 6284402 B1	12						USPAT
US 6171721 B1	10						USPAT

US-PAT-NO: 6171721

DOCUMENT-IDENTIFIER: US 6171721 B1

TITLE: Sputter-deposited fuel cell membranes and electrodes

----- KWIC -----

Detailed Description Text - DPMX (16):

Target 4 may be sputter-deposited onto an anode carrier, a cathode carrier, carbon paper, or other suitable material. The resulting sputter-coated carrier or paper materials are used to fabricate MEAs. Carrier materials useful for fabricating anodes and cathodes and methods for preparing the carrier materials, including methods to alter the wettability of the carriers, are known. To use the methods described herein to prepare an electrode, a suitable carrier such as carbon paper is suitably secured to the substrate holder 8. Once secured, the suitable carrier is sputter-coated using any of the target 4 materials disclosed herein by following the methods described for sputter-coating electrolyte membranes. For example, if an anode is being constructed, the target 4 material should contain catalysts appropriate for the anode such as a platinum-ruthenium alloy. Alternatively, if a cathode surface is being produced, the target 4 material should contain catalysts appropriate for the cathode such as platinum. Other useful target materials include Ni, Ti, Zr, Sn, SnO, sub. 2, Ru, Pt, Os, Ir, Mo, sub. 3, Re, Pd, Mo, Nb, RuO, sub. 2, alloys thereof, and other similar materials.

Current US Class - CLASS (2):

429

United States Patent

Narayanan et al.

(10) Patent No.: US 6,171,721 B1
(45) Date of Patent: Jan. 9, 2001

(54) SPUTTER-DEPOSITED FUEL CELL MEMBRANES AND ELECTRODES

(75) Inventors: Subramanyam B. Narayanan, Alhambra; Barbara Jettles-Narayanan, St. Martin; William Chen, Los Angeles; Ron P. Ruiz, Alhambra; Thomas L. Valdez, Covina, all of CA (US)

(73) Assignee: California Institute of Technology, Pasadena, CA (US)

(*) Notice: Under 35 U.S.C. 154(b), the term of this patent shall be extended for 0 days.

(21) Appl. No.: 09/172,104

(22) Filed: Sep. 22, 1998

Related U.S. Application Data

(60) Provisional application No. 60/099,472, filed on Sep. 22, 1997.

(51) Int. Cl.⁷ H01M 11/00

(52) U.S. Cl. 429/40; 429/42; 429/44; 429/40; 204/290 R; 204/283; 204/192.14; 204/296

(56) Field of Search 204/283, 294, 296, 429/40, 41, 42, 44

References Cited

U.S. PATENT DOCUMENTS

4,275,126 6/1981 Bergman et al. 429/30

4,882,232 11/1989 Bugue et al. 428/613
5,084,144 1/1992 Roddy et al. 204/290 R
5,242,764 9/1993 Deak 429/20
5,277,996 1/1994 Marchetti et al. 429/44
5,572,132 4/1996 Schmitt et al. 204/192.26
5,641,586 6/1997 Wilson 429/30
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* cited by examiner

Primary Examiner—Bruce F. Bell
(74) Attorney, Agent, or Firm—Fish & Richardson P.C.

(57) ABSTRACT

A method for preparing a membrane for use in a fuel cell membrane electrode assembly includes the steps of providing an electrolyte membrane, and sputter-depositing a catalyst onto the electrolyte membrane. The sputter-deposited catalyst may be applied to multiple sides of the electrolyte membrane. A method for forming an electrode for use in a fuel cell membrane electrode assembly includes the steps of obtaining a catalyst, obtaining a backing, and sputter-depositing the catalyst onto the backing. The membranes and electrodes are useful for assembling fuel cells that include an anode electrode, a cathode electrode, a fuel supply, and an electrolyte membrane, wherein the electrolyte membrane includes a sputter-deposited catalyst, and the sputter-deposited catalyst is effective for sustaining a voltage across a membrane electrode assembly in the fuel cell.

61 Claims, 3 Drawing Sheets

Document ID	Pages	U	S	C	P	Kind Codes	Source
US 6326098 B1	11						USPAT
US 6284402 B1	12						USPAT
US 6171721 B1	10						USPAT
US 6162267 A	16						USPAT
US 6143443 A	90						USPAT
US 6131051 A	6						USPAT
US 6127061 A	16						USPAT

US-PAT-NO: 6127061

DOCUMENT-IDENTIFIER: US 6127061 A

TITLE: Catalytic air cathode for air-metal batteries

RWIC

Brief Summary Text - BREF (14):

Thin porous carbon paper based electrodes, such as disclosed in U.S. Pat. No. 3,912,358, solves the bulk problem and has a shortened diffusion path. Unfortunately, thin porous carbon paper substrates are very fragile, and they are subject to excessive flooding with electrolyte which interferes with the access of the gas to the electro catalytic sites of the electrodes. To control the flooding, the carbon papers are often rendered hydrophobic by means of, for example, a Teflon coating which increases their electrical resistivity. In addition, because they are structurally weak, they tend to break in handling, as well as when they operate under moderate gas pressures. Finally, the wet-proofed carbon papers have to be dense to provide a minimum of structural integrity. This characteristic confines a catalytic layer to a surface coating bonded merely to one face of the paper substrate, and being paper, they are inherently nonuniform with respect to porosity. Another thin electrocatalytic gas diffusion electrode comprises a substantially uniform, open pore carbon or graphite substrate, having a thickness in the range to about 5 to 40 mils, and preferably about 10 to about 35 mils and includes a mixture of Teflon or similar wet-proofing particles and catalytic carbon particles imbedded and added within the cloth pores. This type electrode has improved electrochemical performances as well as improve structural strength and is suitable for use in free-flowing electrolytic electrochemical cells. The catalytic carbon particles are either metal-free catalytic carbon particles or finely divided high surface area carbons carrying suitable known noble metal catalytic particles, including platinum, palladium, rhenium, iridium, ruthenium, and silver, depending on the environment (e.g., acid or alkaline, air or hydrogen, and on operating conditions: temperature, current density and intended length of service).

Current US Class - CLAS (1):

429

United States Patent [19]

Shun et al.

Patent Number: 6,127,061
Date of Patent: Oct. 3, 2000

[54] CATALYTIC AIR CATHODE FOR AIR-METAL BATTERIES

[75] Inventors: You-Kuang Shun, Scott Shanghai; Chou-Lai Lou, Shenzhen, both of China

[73] Assignee: High-Density Energy, Inc., Azusa, Calif.

[21] Appl. No.: 09/234,008

[22] Filed: Jan. 26, 1999

[51] Int. Cl.⁷

[52] U.S. Cl. 429/40; 429/42; 429/44; 429/27; 429/59; 429/133; 429/162; 429/163; 429/164; 429/165

[58] Field of Search

429/27, 12, 59, 133, 162, 163, 164, 165; 204/282, 283, 290 R

[56] References Cited

U.S. PATENT DOCUMENTS

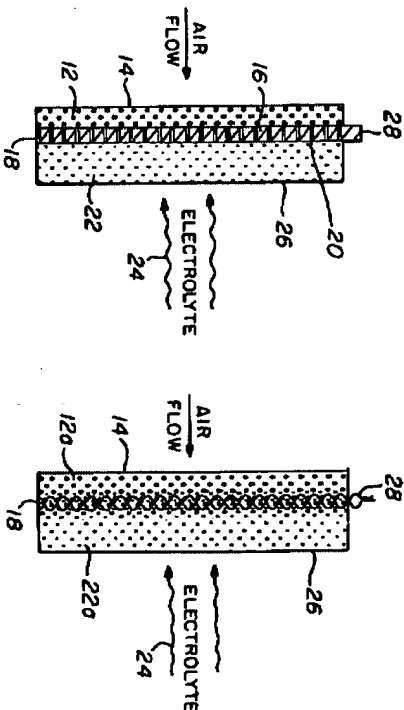
- 3,912,358 10/1973 Davis et al. 136/86 D
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4,302,287 5/1981 Samuels et al. 429/15
4,377,406 3/1981 Solomon 320/423

Primary Examiner—Bruce F. Ball
Attorney, Agent, or Firm—Jones, Day, Reavis & Pogue

ABSTRACT

An air cathode for use in an electrochemical cell or battery having an air permeable and water impermeable layer, an electrically conductive middle layer and a catalytic layer comprising a mixture of carbon particles, particulate materials, having a high surface area, metal hydroxides, and hydrophobic particles.

59 Claims, 6 Drawing Sheets



Document ID	Page	U	S	C	P	Kind Code	Source
22	US 6045938 A	7					USPAT
23	US 6010606 A	9					USPAT
24	US 5928806 A	9					USPAT
25	US 5865968 A	10					USPAT
26	US 5783325 A	14					USPAT
27	US 5729427 A	10					USPAT
28	US 5716437 A	13					USPAT

US-PAT-NO: 5716437

DOCUMENT-IDENTIFIER: US 5716437 A

TITLE: Materials for use in electrode manufacture

----- RWIC -----

Brief Summary Text - BREV (28):

The term "catalyst" will be well understood by a person skilled in the art by meaning a catalyst that when incorporated into a gas diffusion electrode facilitates an electrochemical reaction, for example the catalyst may be selected from the platinum group metals (ie platinum, palladium, rhodium, ruthenium, iridium and osmium), gold, silver or a base metal or base metal oxide, or an alloy or mixture comprising one or more of these metals, preferably supported on a conductive substrate, such as carbon.

Current US Class - CLASS (4):

429

United States Patent (19)

Denton et al

(14) MATERIALS FOR USE IN ELECTRODE MANUFACTURE

(75) Inventor: Jan Denton, Reading; John M. Thompson, Reading, all of Great Britain

(73) Assignee: Johnson Matthey Public Limited Company, London, England

(21) Appl. No.: 613,387

(22) Filed: Mar 7, 1996

(30) Foreign Application Priority Data

Mar 9, 1995 (GB) United Kingdom 9506713

(51) Int. Cl. C C09D 11/00

(52) U.S. Cl. 204/283; 204/286; 204/290 B; 204/291; 204/292; 204/293; 429/40; 429/42; 429/44; 429/45

(58) Field of Search 204/282, 283, 286, 290 B, 291, 292, 293; 429/40, 42, 44, 45

(56) References Cited

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5,346,780 9/1994 Szabo 429/42

(11) Patent Number: 5,716,437
(45) Date of Patent: Feb. 10, 1998

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0 305 545 3/1989 European Pat. Off.
0 309 337 3/1989 European Pat. Off.
0 360 062A2 11/1993 European Pat. Off.
0 622 861A1 11/1994 European Pat. Off.
1 285 859 8/1972 United Kingdom.
WO 94/25993 11/1994 WIPO.

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Jan D. Reisterick, "Modified Gas Diffusion Electrodes for Proton Exchange Membrane Fuel Cells", Extended Abstracts, vol. 86-1, May 4-9, 1986, p. 660.
Pearl Abstracts of Japan, JP 58147575 published Feb. 9, 1983.

Primary Examiner—Bruce F. Bell
Attorney Agent, or Firm—Cushman Dabry & Cushman IP Group, Philadelphia and Suro LLP

(57) ABSTRACT

An improved ink material, particularly for use in printing processes and its use in improved manufacturing processes for higher performance electrodes for application in fuel cells and other electrochemical devices is disclosed.

17 Claims, 5 Drawing Sheets

United States Patent	[19]	Patent Number:	5,865,881
Mori et al		Date of Patent:	Feb. 2, 1999
[54] ELECTROLESS PLATING BATH OF IRIDIUM			
[75] Inventors: Hiroaki Mori, Toyonaka, Shoji Mazumura, Kawasaki; Katsuhiko Oguro, Izumi-Shi; Eiichi Torihara, Yto, all of Japan			
	3928634 A1	2/1991	Germany.
	58-197381	11/1983	Japan.
	2,20709	8/1985	Japan.
	60-128780	8/1985	Japan.
	60-162780	8/1985	Japan.

6 Claims, 2 Drawing Sheets

